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Isolation and characterization of a N. CRASSA silencing geneand uses thereof

The present invention relates to the isolation and characterization of a Neurospora crassa gene encoding for an essential activity in the co-suppression process and to uses and applications thereof in vegetal, animal and fungine fields.

The production of transgenic organisms is of large utility both in basic and applied biological research. The transgenic DNA is usually integrated in the genome and transferred as a Mendelian character. However, in various instances, the transgene introduction induces gene silencing phenomena (Flavell, R.B. 1994), i.e. the repression of the expression of the transgene itself and/or of one or more endogenous homologous genes.

The gene silencing (suppression of gene expression) can act at two levels: transcriptional (transinactivation) where transgenes contain sequences homologous to the silenced gene promoter (Vaucheret, 1993); and post-transcriptional (co-suppression) which requires homologies between coding regions (Flavell, 1994; Stam et al., 1997; Baulcombe, 1996).

Generally the silencing induced by a transgene requires an almost complete sequence homology (from 70% to 100%) between transgene and silenced gene sequences (Elkind, 1990).

In the Neurospora crassa filamentous fungus, during the vegetative phase, the presence of transgenes induces a post-transcriptional gene silencing phenomenon, named "quelling" (Cogoni et al., 1996).

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By using the al-1 gene (albino 1) (Schmidhauser et al., 1990) as silencing visual marker, many features of the phenomenon have been discovered (Cogoni et al., al-1 gene "quelling" Particularly the 1) the gene Neurospora is characterized in that: silencing is reversible further to the loss of transgene copies; 2) the reduction of mRNA basal level results from a post-transcriptional effect; 3) transgenes containing at least a region of 132 base pairs which is identical to the region encoding for the target gene are sufficient to induce the "quelling"; 4) the duplication of promoter sequences is ineffective to induce the silencing; 5) the "quelling" exhibits a dominant behavior in eterocarions containing both transgenic and untransformed nuclei, indicating the involvement of a trans-acting diffusible molecule among the nuclei; 6) the expression of aberrant RNA transcribed by the transgenic locus strictly correlated to silencing, suggesting that the "quelling" can be induced and/or mediated by a transgenic RNA molecule.

Therefore homologies between *Neurospora* silencing and plant co-suppression can be pointed out. The gene silencing in *Neurospora* is reversible, as result of transgenic copies instability during mitotic phase; in plants also the co-suppression reversion is associated with the reduction of transgene copy number, resulting from intra-chromosomal recombination during mitosis or meiosis (Mittelstein Scheid et al., 1994; Stam et al., 1997). Thus both in plants and in *Neurospora* the transgene presence is required to maintain the silencing. As in *Neurospora*, a decrease of the mRNA basal level of the silenced gene results from a post-transcriptional

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mechanism (Dehio and Schell 1994; van Blokand et al., 1994; de Carvalho et al., 1995). Furthermore to induce the "quelling", transgenes must contain a portion of the silencing target gene coding sequence, being the promoter region ineffective. In plants coding regions with no promoter sequences can induce silencing (van Blokand et al., 1994) and, as in the "quelling", promoters or functionally active gene products are not required for the co-suppression.

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One of the similarities between "quelling" and cosuppression in plants is that both mechanisms diffusion In Neurospora mediated by factors. eterokaryotic strains, nuclei wherein the albino-1 gene is silenced are able to induce the al-1 gene silencing of the other not transformed nuclei, all sharing the same cytoplasmic environment (Cogoni et al., 1996). In plants the presence of a diffusion factor results from the fact that the co-suppression is effective in inhibiting the replication of Tobacco Etch Virus (TEV), a RNA virus with an exclusively cytoplasmic cycle. The occurrence of highly diffusible factors, which are effective to mediate the co-suppression, has been demonstrated using the grafting technique in tobacco (Palaqui et al., 1997), showing that silenced tobacco plants are able to transfer the silencing to non-silenced plants through grafting.

The fact that "quelling" and co-suppression share all these features suggests that mechanisms involved in post-transcriptional gene silencing in plants and in fungi can be evolved by an ancestral common mechanism.

Recently gene inactivation phenomena resulting from transgene introduction have been disclosed in animals. In Drosophila melanogaster the location of a transgene close

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to heterochromatic centers results in a variegate expression (Wallrath and Elgin, 1995; Pirrotta, V., 1997). Similar expression profiles have been observed when the reference transgene is within tandem arrayed transposons, indicating that tandem repeats are effective to induce the chromatin condensation. (Dorer and Henikoff, 1994). Again in *Drosophila* Pal-Bhadra et al. (1997) have observed that the transgene introduction can lead to gene inactivation phenomena, similar to the cosuppression.

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Gene silencing phenomena resulting from transegene sequence repeats have been disclosed recently in mammalians.

Garrick et al. (1998) produced mouse transgenic lines wherein 100 transgenic copies are present in a unique locus and are repeats-arrayed in direct tandem. The transgene expression has been disclosed to be inversely proportional to the number of occurring copies, indicating that silencing phenomena dependent on repeat copies are present also in mammalians.

It has been recently found that double stranded RNA molecules can induce a sequence-specific silencing in several organisms (Fire A., 1999). The mechanism known as dsRNAi (double stranded RNA interference) acts at a post-transcriptional level by inducing sequence-specific degradation of homologous mRNAs (Montgomery, Xu and Fire, 1998). Under this aspect, dsRNAi and quelling in Neurospora are similar mechanisms, both of them acting at a post-transcriptional level. In addition, both RNA-induced silencing and DNA-induced silencing can be transmitted from cell to cell.

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Therefore the identification of *Neurospora* genes which are involved in the silencing is the first step to modulate the same process in plants, animals and fungi. The silencing modulation is of great relevance when transgenic organisms able to express the desired phenotype are produced.

The authors of the present invention have already isolated Neurospora crassa strains mutated at essential functions for gene silencing (Cogoni and Macino, 1997); 15 independent isolated mutants define three complementation groups, thus identifying the qde-1, qde-2 and qde-3 genes (qde stands for "quelling"-deficient), whose products are essential to the silencing machinery. qde genes are essential to the Neurospora silencing, as suggested by the fact that silencing of three independent genes (al-1, al-2 and qa-2) is impaired by qde mutations (Cogoni and Macino, 1997).

The authors of the present invention have already identified qde-3 gene (PCT WO 00/327885) and qde-1 gene (PCT WO 00/50581).

The authors of the invention have identified and cloned now one out of Neurospora qde genes, the qde-2 gene, thus identifying one of required factors for silencing. By considering the similarity between "quelling" and co-suppression, genes orthologous to the isolated gene are involved in co-suppression and more generally in gene silencing in other organisms, like plants, fungi and animals.

The present invention can be applied with reference to two general scopes: 1) silencing potentiation as a tool for inactivating more effectively and durably a

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desired gene, and 2) silencing suppression to obtain a better expression of the introduced transgenes.

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The isolated qde-2 gene can be introduced alone or with qde-1 and/or qde-3 genes in plants, animals or fungi, in order to inactivate the expression of selected genes. The aim is to activate a sequence-specific silencing mechanism both in deficient organisms and in organisms wherein the same is not very efficient. The gene silencing can be induced also by introducing specific double stranded DNA or RNA sequences, homologous to the gene to be inactivated.

As to the silencing potentiation, the overexpression of one or more genes controlling phenomenon can lead to higher efficiency and/or stability thereof. Therefore the introduction of qde-2 gene or of homologous genes thereof in organisms can constitute a gene to repress more effectively functions. Particularly this approach is specially useful in plants wherein the co-suppression is usually used for the "knock-out" of gene functions. In plants again the gene silencing potentiation can be used to obtain lines resistant to pathogen virus, by introducing transgenes encoding for viral sequences, in order to achieve the expression inhibition of the virus itself (Flavell et al., 1994).

Analogous applications are suitable for animals, wherein some indications suggest that silencing can inhibit the suitable expression of introduced transgenes (Garrick et al., 1998).

On the contrary, there are instances wherein it is desirable not to have or to reduce the gene silencing, i.e. where a transgene is to be over-expressed. It is

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known that the co-suppression is strictly correlated both with the presence of an high copy number of transgene, and with a transgene high expression. This correlation can hamper the production of transgenic. organisms which express a transgene at high levels, because more high is the expression and/or the copy number, more probable is to evoke silencing responses. As above mentioned, analogous mechanisms of inactivation, dependent on a high copy number, have been disclosed in animals. In these circumstances plant or animal lines, totally or partially ineffective silencing, constitute an ideal recipient wherein the desired gene can be over-expressed. The invention can be applied within this scope using different approaches:

A) Identification and production of mutant lines in genes homologous to qde-2 gene, in plants, animals and fungi.

The identification of Neurospora qde-2 gene, essential for silencing mechanism, can allow the isolation of mutant lines in other organisms, mutated in genes homologous to qde-2. For example by means of amplifications using degenerated primers, designed from the most conserved regions of qde-2 gene, mutant lines in homologous genes can be identified, by analysis of insertion mutant gene banks, already available for many plant species. Both in fungi and animals such mutants can be obtained, following the identification of the homologous gene, by means of "gene disruption" techniques using homologous recombination.

B) Reduction of qde-2 gene expression

Other strategies for the production of silencing-deficient lines comprise the use of Neurospora qde-2 gene

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or homologous genes thereof. qde-2 or homologous genes can be introduced into suitable expression vectors to express them in an anti-sense orientation in order to inhibit the expression of resident endogenous genes. Alternatively portions of qde-2 or of homologous genes can be over-expressed, in order to obtain a negative dominant effect and thus blocking the function of qde-2 endogenous genes.

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The authors of the present invention have cloned and characterised the *Neurospora crassa qde-2 gene*. The sequence analysis of the *qde-2* gene detected a region having a significant homology with the sequence of a *C*. elegans gene, rde-1, involved in the dsRNA mediated interference (Tabara et al., 1999).

The authors of the invention for the first time have demonstrated that the transgene induced posttranscriptional gene silencing and the dsRNA interference share common genetic mechanisms. This supports hypothesis that the sequence specific gene silencing phenomena evolved from an ancestral mechanism aimed to protect the genome against transposons. Furthermore, the results of the authors suggest that dsRNA molecules are involved in the post-transcriptional gene silencing in fungi. dsRNA molecules could be produced directly from integrated trangenes as a result of the presence of inverted repeats or as an out come of transcription from convergent inverted promoters. Alternatively, single stranded aberrant RNA may be used as a template by an RNA-dependent RNA polymerase (such as QDE-1 protein) able to produce dsRNAs.

Within the scope of the invention the term homology is intended as similarity, i.e. number of identical

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residues + number of conserved residues with respect to the total residues of the considered sequence.

Therefore it is an object of the present invention an isolated nucleic acid molecule encoding for a protein characterized in having a silencing activity and in comprising a domain responsible for dsRNA interference, wherein the domain is at least 25% homologous with the amino acid sequence from aa. 373 to aa. 910 of sequence in fig. 1 (SEQ ID No. 2). Preferably the domain is at least 30% homologous with the amino acid sequence from aa. 373 to aa. 910 of sequence in fig. 1 (SEQ ID No. 2). More preferably the domain is at least 38% homologous with the amino acid sequence from aa. 373 to aa. 910 of sequence in fig. 1 (SEQ ID No. 2). Most preferably the domain comprises the amino acid sequence from aa. 373 to aa. 910 of sequence in fig. 1 (SEQ ID No. 2). According to a particular embodiment the isolated nucleic acid molecule encodes for a protein having the amino acid sequence of fig. 1 (SEQ ID No. 2) or functional portions thereof. Even more preferably the isolated nucleic acid molecule has the sequence of fig. 1 (SEQ ID No. 1) or its complementary sequence.

A further object of the invention is an expression vector comprising, under the control of a promoter which directs the expression in bacteria, the isolated nucleic acid molecule of the invention. Those skilled in the art will appreciate that any plasmid suitable for a correct and effective expression of the protein of the expression in bacteria can be used and it is within the scope of the invention.

A further object of the invention is an expression vector comprising, under the control of a promoter which

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directs the expression in plants or in specific plant organs, the isolated nucleic acid molecule of invention, both in a sense and anti-sense orientation. Those skilled in the art will appreciate that any plasmid suitable for a correct and effective expression of the protein of the invention in plants or in specific plant organs can be used and it is within the scope of the invention.

A further object of the invention is an expression 10 vector comprising, under the control of a promoter which directs the expression in fungi, the isolated nucleic acid molecule of the invention, both in a sense and antisense orientation. Those skilled in the art will appreciate that any plasmid suitable for a correct and effective expression of the inventive protein in fungi can be used and it is within the scope of the invention.

A further object of the invention is an expression vector comprising, under the control of a promoter which directs the expression in animals, the isolated nucleic acid molecule of the invention, both in a sense and antisense orientation. Those skilled in the art will appreciate that any plasmid suitable for a correct and effective expression of the protein of the invention in animals can be used and it is within the scope of the invention.

A further object of the invention is a prokaryotic organism transformed by using the expression vector active in bacteria of the invention.

A further object of the invention is a plant or a specific plant organ transformed by using the expression vector active in plants of the invention.

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A further object of the invention is a plant mutated at the isolated nucleic acid molecule of the invention having a reduced or inhibited silencing activity.

A further object of the invention is a fungus transformed with the expression vector of the invention active in fungi.

A further object of the invention is a fungus mutated at the isolated nucleic acid molecule of the invention and having reduced or inhibited silencing activity.

A further object of the invention is a non-human animal transformed with the expression vector of the invention active in animals.

A further object of the invention is a non-human animal mutated at the isolated nucleic acid molecule of the invention and having a reduced or inhibited silencing activity.

A further object of the invention refers to a protein characterized in having a silencing activity and in comprising a domain responsible for dsRNA interference, wherein the domain is at least 25% homologous with the amino acid sequence from aa. 373 to aa. 910 in fig. 1 (SEQ ID No. 2). Preferably the domain is at least 30% homologous with the amino acid sequence from aa. 373 to aa. 910 in fig. 1 (SEQ ID No. 2). More preferably the domain is at least 38% homologous with the amino acid sequence from aa. 373 to aa. 910 in fig. 1 (SEQ ID No. 2). Most preferably the domain comprises the amino acid sequence from aa. 373 to aa. 910 in fig. 1 (SEQ ID No. 2). According to a particular embodiment the isolated nucleic acid molecule encodes for a protein

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having the amino acid sequence of fig. 1 (SEQ ID No. 2) or functional portions thereof.

It is within the scope of the present invention the use of the isolated nucleic acid molecule of the invention to modulate gene silencing in plants, animals and fungi.

The present invention now will be described by way of non limiting examples with reference to the following figures:

Figure 1: The isolated nucleic acid molecule of the 5.7 Kb fragment containing the qde-2 gene and flanking sequences (SEQ ID No.1). The amino acid sequence (SEQ ID No. 2) is shown above the nucleotide sequence.

Figure 2: It is schematically represented the pMXY2 plasmid insertion site, in the 80 mutant, used for insertional mutagenesis and consequent polimorphism of the restriction fragments by mean of DNA southern blot of a WT strain and of 80 and 820 mutant strains by using the entire restored flanking region as probe. The 820 mutant has a complete deletion of the qde-2 gene.

Figure 3: Multiple alignment, at the conserved region, among qde-2 and other proteins belonging to ago-elF2C family: A. thaliana ago-1; rabbit elF2C; C. elegans rde-1. Identical amino acids are shown in bold.

25 MATERIALS AND METHODS

E. coli strains

E. coli strain HB101 (F, hsdS20(rb, mb), supE44,
recA13, ara14, proA2, rspL20(str, xyl-5) was used for
cloning.

30 Neurospora crassa strains and growing conditions

Neurospora crassa following strains, supplied by Fungal Genetic Stock Center (FGSC, Dpt. Of Microbiology,

WO 01/53475

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University of Kansas Medical Ctr. Kansas City, KA) were used:

- Wild type (FGSC 987);
- qa-2/aro9 (FGSC 3957A), (FGSC 3958a).

The 6XW strain (Cogoni et al., 1996) was obtained upon transformation of the FGCS 3958a strain with pX16 plasmid (Cogoni et al., 1996). This plasmid contains the qa-2 gene used as selective marker and the al-1 coding sequence.

The mutant strains M7, M20 (qde-1); M10, M11 (qde-2); M17, M18 (qde-3) are described in Cogoni and Macino, 1997.

The qde mutants were obtained by UV mutagenesis. As recipient the transforming strain (6xw) silenced at the albino-1 gene was used. qde mutants were selected for their ability to recover a wild type unsilenced phenotype and then classified in three different complementation groups. By analyzing the al-2 gene quelling frequency all of qde used mutants are defective for the general silencing mechanism.

Complementation assays with not forced heterocaryons were carried out according to Davis and DeSerres, 1970.

Plasmids and libraries

25 The plasmid pMXY2, disclosed in Campbell et al. 1994, used for insertional mutagenesis was obtained from Fungal Genetic Stock Center (FGSC, Dpt. Of Microbiology, University of Kansas Medical Ctr. Kansas City, KA). The plasmid contains the *Bm1* gene (allele responsible of the benilate drug resistance), that was used as selective marker after transformation. The genomic DNA containing

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the qde-2 gene was isolated from a N. Crassa gene library in cosmids. (Cabibbo et al., 1991).

N. crassa transformation

Spheroplasts were prepared according to the Akins and Lambowitz (1985) protocol.

Southern Blot Analysis

Chromosomal DNA was prepared as disclosed by Irelan et al., 1993. 5 μg of genomic DNA were digested and blotted as reported in Maniatis et al.

DNA probes were: a) as to the al-1 gene the probe is represented by a XbaI-ClaI restriction fragment of pX16 (Cogoni et al., 1996); b) as to the BmI gene the probe is represented by the 2.6Kb SalI fragment of pMXY2.

Northern Blot Analysis

N. crassa total RNA was extracted according to the protocol described by Cogoni et al., 1996. The mycelium was grown for two days at 30°C, then powdered in liquid nitrogen before RNA extraction. For Northern analysis 10 μ g of RNA were formaldehyde denatured, electrophoresed on a 1% agarose, 7% formaldehyde gel, and blotted over Hybond N (Amersham) membranes. Hybridization was carried out in 50% formamide in the presence of ³²P labeled DNA probe 1.5x10⁶ cpm/ml.

RESULTS

25 Isolation of silencing mutant by insertional mutagenesis

Previously a Neurospora strain (6XW) wherein the albino-1 resident gene was steadily silenced was used for UV mutagenisis that brought to the isolation of qde ("quelling" deficient) mutants in N. crassa induced gene silencing (Cogoni and Mancino 1997).

The 6XW strain shows an albino phenotype due to the lack of carotenoid biosynthesis, as results by the

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silencing of the albino 1 gene expression (Schmidhauser et al., 1990). A mutation interfering with the silencing machinery is easily detectable by producing a wild type phenotype (bright orange) of the carotenoid biosynthesis. 5 By means of complementation assays it was possible to establish that qde mutants belong to three complementation groups, indicating the presence of three genetic loci involved in the Neurospora silencing mechanism. In order to isolate the qde genes 10 insertional mutagenesis was carried out with the 6XW previously used for UV mutagenesis. The insertional mutagenesis was carried out by transforming the 6XW strain with a plasmid, taking advantage of the fact that, after the transformation, plasmids are 15 randomly inserted in the Neurospora crassa genome. The mutagenesis was carried out transforming the 6XW silenced strain with pMXY2 (see Materials and Methods) which contains the benilate resistance as selective marker. Transformed strains able to grow in the presence of 20 benilate containing medium and showing a wild type phenotype for the carotenoid biosynthesis were selected. Out of 50.000 isolated independent transformed strains, a benilate resistant strain (80) was isolated, which showed the bright orange phenotype expected for a qde gene mutation. In order to verify that the silencing release 25 was effectively due to a qde gene mutation and not to the loss of al-1 transgene copies, the genomic DNA of the strain 80 was extracted and digested with SmaI and HindIII restriction enzymes. After blotting, 30 hybridized with a probe corresponding to the coding sequence of al-1. The SmaI site is present only once in the al-1 transgene containing plasmid and the digestion

by using said enzyme produces a 5.5Kb fragment corresponding to tandem arrayed al-1 transgenes, while a 3.1Kb fragment is expected from the resident al-1 locus. The number of al-1 transgenic copies present in the 80 strain is comparable to that present in the silenced 6XW strain.

The strain 80 is mutated in qde-2 gene

The strain 80 was assayed in a heterokaryon assay with a wild type strain and with M7, M20 (qde-1) M10, M11 (qde-2), M17, M18 (qde-3) mutants and with a wild strain (Cogoni and Macino, 1997). As shown in Table 1 the al-1 gene silencing is restored producing an albino phenotype in all of heterocaryons but M10 and M11. This behavior is consistent with the presence of a qde-2 gene recessive mutation in the strain 80.

Table 1
Reciprocal heterokaryons among the mutant 80 and previously characterized qde mutants.

	80	M7	M20	M10	M11	M17	M18
80	WT	AL	AL	WT	WT	AL	AL
M7		WT	WT	AL	AL	AL	AL
м20			WT	AL	AL	AL	AL
м10				WT	WT	AL	AL
M11					WT	AL	AL
М17						WT	WT
M18							WT

WT = heterokaryon with a wild type phenotype for

20 carotenoid accumulation;

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AL = heterokaryon with an albino phenotype wherein the al-1 gene silencing is restored.

Recovery of sequences flanking the pMXY2 plasmid integration site

WO 01/53475

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In order to recover sequences flanking the integration site or sites the following methodology was carried out. The genomic DNA of strain 80 was digested with Aat II enzyme. Subsequently the genomic DNA was ligated and the product used to transform *E. coli* cells that was screened in an ampicillin-containing medium. PQc1 plasmid was recovered and a DNA fragment containing sequences flanking the integration site was isolated from it by using Aat II and Cla I enzymes.

Isolation of genomic clones, their subcloning and complementation of the qde-2 mutant

The fragment from pQc1 plasmid was used to probe a Neurospora crassa genomic library in cosmids. Three cosmids 6G10, 20C1 and 23F2 containing about 35 Kb genomic DNA inserts, were isolated. Such cosmids were used in transformation experiments of M11 and 80 mutants. All of cosmids are able to restore the al-1 gene silencing in the two mutants, determining the appearance of an albino phenotype. The 20C1 cosmid was used to subclone a 5.7 Kb BamHI-BamHI fragment. This subclone was used for transformation experiments and resulted to be able to complement the qde-2 phenotype, indicating that a qde-2 functional gene is present in this plasmid.

Isolation and sequence of the qde-2 cDNA

The sequence of BamHI-BamHI region allowed to deduce the amino acid sequence of the QDE-2 protein. The qde-2 gene encodes for a 938 aa. putative protein (104 KDa). The genomic clone does not contain any introns since the reading frame does not contain any interruptions and intron acceptor and donor sequences were not identified (Fig. 1, Seq. ID No 1, 2).

WO 01/53475

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The qde-2 gene comprises an homologous domain with encoding genes for proteins that are responsible for dsRNA interference

The 938 aa sequence (SEQ ID No. 2) was used to search in database of amino acid sequences, by using the BLASTP algorithm. As showed in fig. 3, the search identified significant homologies with argonaute-1 gene [with expected values (E value) of 2e-57] of A. Thaliana (mutants of this gene show developmental anomalies); rde-1 gene [with expected values (E value) of 1e-23] of C. elegans, involved in gene silencing phenomena induced by double stranded RNA; elF2C gene [with expected values (E value) of 5e-60] of rabbit isolated as an element belonging to transcription beginning complex.

15 Plant expression vector

The ade-2 gene was inserted, in a sense orientation, into a vector containing a plant expression "cassette", including the 35S promoter and the PI-II "terminator" sequences. The vector also includes the Streptomyces hygroscopicus bar gene, which confers the phosphinotricine herbicide resistance to transformed plants. In an analogous vector to the above mentioned one, qde-2 was inserted in an anti-sense orientation with respect to the 35S promoter.

The obtained vectors can be utilized to over-express the qde-2 gene in plants, or to repress the gene expression of resident genes, which are homologous to qde-2.

Fungus expression vector

The qde-2 gene was inserted in a vector containing a fungal specific expression "cassette", comprising the A. nidulans trpC gene promoter and terminator, both in a

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sense and an anti-sense orientation. In addition the vector contains the bacterial hph gene, which confers the hygromicine drug resistance. The sense plasmid can be used to over express the qde-2 gene, whereas the antisense plasmid is used to repress the expression of qde-2 homologous genes in various fungine species.

Mammalian expression vector

The qde-2 gene was inserted in a vector containing a mammalian specific expression "cassette", including the cytomegalovirus (CMV) promoter and SV40 termination and polyadenylation sequences both in a sense and anti-sense orientation. The vector includes also the neomicine phototransferase gene, as marker for mammalian cell selection. The sense plasmid can be used to over express the qde-2 gene, whereas the anti-sense plasmid can be used to repress the expression of qde-2 homologous genes in various mammalian species.

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21

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WO 01/53475

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Claims

- r. An isolated nucleic acid molecule encoding for a protein characterized in having a silencing activity and in comprising a domain responsible for dsRNA interference, wherein the domain is at least 25% homologous with the amino acid sequence from aa. 373 to aa. 910 of SEQ ID No. 2.
- 2. An isolated nucleic acid molecule encoding for a protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 1, wherein the domain is at least 30% homologous with the amino acid sequence from aa. 373 to aa. 910 of SEO ID No. 2.
 - 3. An isolated nucleic acid molecule encoding for a protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 2, wherein the domain is at least 38% homologous with the amino acid sequence from aa. 373 to aa. 910 of SEQ ID No. 2.
 - 4. An isolated nucleic acid molecule encoding for a protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 3, wherein the domain is the amino acid sequence from aa. 373 to aa. 910 of SEQ ID No. 2.
 - 5. An isolated nucleic acid molecule encoding for a protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 4, wherein said isolated nucleic acid molecule encodes for a protein having the amino acid sequence of SEQ ID No. 2, or functional portions thereof.

complementary sequence.

WO 01/53475

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6. An isolated nucleic acid molecule encoding for a protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 5, wherein said isolated nucleic acid molecule has the sequence of SEQ ID No. 1 or its

23

- 7. Expression vector comprising, under the control of a promoter that directs the expression in bacteria, the isolated nucleic acid molecule according to any one of claims 1-6.
- 8. Expression vector comprising, under the control of a promoter that directs the expression in plants or in specific plant organs, the isolated nucleic acid molecule according to any one of claims 1-6, both in a sense and anti-sense orientation.
- 9. Expression vector comprising, under the control of a promoter that directs the expression in fungi, the isolated nucleic acid molecule according to any one of claims 1-6 both in a sense and anti-sense orientation.
- 10. Expression vector comprising, under the control of a promoter that directs the expression in animals, the isolated nucleic acid molecule according to any one of claims 1-6 both in a sense and anti-sense orientation.
- 11. Prokaryotic organism transformed by using the expression vector active in bacteria according to claim 7.
- 12. Plants or a specific plant organ transformed by using the expression vector active in plants according to claim 8.
- 13. Plant mutated at the isolated nucleic acid molecule according to any one of claims 1-6 having a reduced or inhibited silencing activity.

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WO 01/53475

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14. Fungus transformed by using the expression vector active in fungi according to claim 9.

24

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- 15. Fungus mutated at the isolated nucleic acid molecule according to any one of claims 1-6 having a reduced or inhibited silencing activity.
- 16. Non-human animal transformed by using the expression vector active in animals according to claim 10.
- 17. Non-human animal mutated at the isolated nucleic acid molecule according to any one of claims 1-6 having a reduced or inhibited silencing activity.
 - 18. Protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference wherein the domain is at least 25% homologous to the amino acid sequence from aa. 373 to aa. 910 of SEQ ID No. 2.
 - 19. Protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 18 wherein the domain is at least 30% homologous to the amino acid sequence from aa. 373 to aa. 910 of SEQ ID No. 2.
 - 20. Protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 19 wherein the domain is at least 38% homologous to the amino acid sequence from aa. 373 to aa. 910 of SEQ ID No. 2.
 - 21. Protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 20 wherein the domain is the amino acid sequence from aa. 373 to aa. 910 of SEQ ID No. 2.

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- 22. Protein characterized in having a silencing activity and comprising a domain responsible for dsRNA interference according to claim 21 comprising the amino acid sequence of SEQ ID No. 2 or functional portions thereof.
- 23. Use of the isolated nucleic acid molecule according to any one of claims 1-6 to modulate the gene silencing in plants, animals and fungi.

4/7

Length of cBAMqde2.txt: 5746 bp; Listed from: 1 to: 5746; Translated from: 1039 to: 3852 (ORFs); Genetic Code used: Universal; Lun, 27 ago 1956 18:50

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FIG. 1-1

1284 1293 1302

1266

1275

2/7

1332 1341 1350 PSSNQNLPSKPQTWVVKV TGG ACC GAG CCG AGT TCC AAC CAA AAC CTG CCC AGC AAG CCC CAG ACT TGG GTG GTC AAG GTG GAG E T C D F G K V L N E L T T L D P K GAG AGT GTC GAA ACC TGC GAT TTC GGC AAG GTG CTG AAC GAG CTC ACG ACA CTT GAT CCC AAG CTC 1473 1482 YNVELDALNTIVTHH GAC GGA GAC TTT CCC AAG TAC AAT GTG GAG CTC GAT GCC CTC AAC ACC ATT GTG ACT CAT GCC R A D D N V A V V G R G R F F A I G D D L I CGC GCC GAC GAC AAT GTT GCG GTG GGA AGG GGA AGG TTT TTT GCC ATT GGT GAT GAC CTC ATT V R P H D S P L V I L R G Y F A S V GAA CAA GTG CGG CCC CAT GAC TCC CCT TTG GTC ATC TTG CGA GGA TAT TTT GCC AGC GTC CGA 1.662 A T G R L L L N T N I T H G V F R P G V K L GCT ACC GGA AGA CTT TTA CTC AAT ACC AAC ATC ACG CAT GGT GTC TTC CGT CCT GGG GTC AAA CTT A Q L F Q E L G L D V M D K C N A W N E V T GCA CAG CTG TTT CAG GAA CTT GGA CTT GAC GTA ATG GAC AAA TGC AAT GCC TGG AAC GAA GTA ACC K N Q L N D K M R R V H K V L A K G R V E L AAA AAT CAG CTC AAC GAC AAG ATG CGC AGA GTT CAC AAG GTC CTG GCT AAG GGC CGT GTC GAG TTG F L I D G K I V Y K K C Y R T L N G AAT GCC CCA TTC CTT ATT GAT GGA AAG ATT GTT TAT AAA AAA TGT TAC CGC ACG CTC AAT GGC ATT 1935 1944 1953 DERGKQKDGKEVRY GCT AAC CGT GGC GAC GAA AGG GGG AAG CAA AAG GAT GGT AAA GAA GTC CGA TAT CCG CCC TTG TTC G I P G V Q V G G P T S C Q F Y L R A R E T GGG ATT CCG GGT GTC CAG GTT GGC CCG ACC TCT TGT CAG TTC TAC TTG CGT GCG CGA GAG ACA AAG GAT GGC GCT GCC CCT CCT CCG ACT CCC GGC CTG CCG AGC AAC GCG TAC ATC ACG GTA GCG AAC LTANEADNMIKFACRAPS CTG ACA GCC AAC GAG GCG GAC AAC ATG ATT AAG TTT GCT TGC AGA GCT CCT TCG CTG AAC GCT CAG S I V T K G R O T L G L D K S L T L G K TCT ATC GTG ACG AAA GGC AGA CAG ACA CTT GGT CTT GAT AAA AGC CTG ACG CTT GGC AAG TTC AAG D K E L I T V V G R E L K P P M L T GTT TCG ATC GAC AAG GAG CTG ATC ACC GTT GTC GGG CGT GAG CTC AAG CCT CCG ATG CTT ACG TAC

3/7

S G N K T V E P Q D G G W L M K F V K V A R AGG GGT AAC AAG ACG GTA GAG CCG CAG GAC GGC GGG TGG TTG ATG AAG TTT GTC AAG GTC GCC AGA 2511 2520 2529 2538 2547 2556 P C R K I E K W T Y L E L K G S K A N E G V CCT TGC CGC AAG ATT GAG AAG TGG ACA TAC TTG GAA CTG AAG GGT TCC AAG GCA AAC GAA GGG GTG A M T A F A E F L N R T G I P I N P R F CCG CAA GCT ATG ACC GCT TTT GCC GAA TTC TTG AAC AGA ACG GGC ATC CCG ATT AAC CCC AGG TTC S P G M S M S V P G S E K E F F A K V K E L TCG CCG GGC ATG AGG TC CCA GGG AGC GAA AAA GAG TTC TTT GCC AAA GTG AAG GAA CTC M S S H Q F V V V L L P R K D V A I Y N M ATG AGC TCG CAC CAA TTT GTG GTG GTT CTT TTA CCC AGA AAG GAT GTT GCG ATC TAC AAT ATG GTG K R A A D I T F G V H T V C C V A E K F L S AAG CGG GCT GCC GAT ATC ACA TTT GGC GTT CAC ACA GTC TGT TGT GTA GCC GAA AAG TTC CTT AGC K G Q L G Y F A N V G L K V N L K ACT AAG GGG CAG CTG GGG TAT TTT GCC AAC GTC GGC CTC AAG GTC AAC CTC AAG TTT GGC GGC ACC H P T N L A A G Q S P A S A P S I V G GTC ACC CAT CCG ACC AAT CTA GCG GCT GGA CAA TCG CCT GCA TCG GCT CCC AGT ATT GTC GGC CTG V S T I D Q H L G Q W P A M V W N N P H G O GTC TCA ACC ATC GAC CAA CAC CTT GGA CAA TGG CCT GCA ATG GTT TGG AAC AAC CCG CAC GGC CAG E S M T E Q F T D K F K T R L E L W R S N P GAG TCC ATG ACG GAA CAG TTT ACG GAC AAG TTC AAG ACG CGT CTG GAA CTA TGG CGC AGC AAT CCC R S L P E N I L I F R D G V S N GCA AAC AAC CGC AGT CTC CCC GAG AAT ATC CTG ATT TTC CGC GAT GGC GTC TCC GAG GGA CAG TTC Q M V I K D E L P L V R A A C K L V Y P A CAG ATG GTC ATC AAG GAC GAG CTA CCC CTG GTT CGC GCC GCC TGC AAG CTG GTG TAT CCA GCT GGC 3321 3330 3339 3348 RITLIVSVKRH AAG CTA CCG CGT ATT ACG CTG ATT GTC TCT GTC AAG CGC CAC CAG ACT CGC TTC TTC CCA ACG GAC CAC TAC ACA GTT CTG GTG GAT GAG ATT TTC AGG GCC GAC TAT GGA AAC AAG GCG GCC GAC ACG CTG

4/7

3633	3642	3651	3660	3669	3678	3687
P A Y Y CCT GCG TAC TAT 3699		·		I H Q ATC CAT CAG 3735	K E L AAG GAG CTC 3744	F D A TTT GAC GCC 3753
L D E N CTC GAT GAA AAC 3765	D S V K GAT AGC GTT A 3774			R W G AGA TGG GGT 3801	N S G AAC TCC GGG 3810	A V H GCT GTT CAT 3819
P N L R CCC AAC CTT AGG 3831	N S M Y AAC TCC ATG T 3840		TAG GCT TGT 3858	CAA TTG TGT 3867	GCT GGA ATG 3876	TAC TGG AGC 3885
ATA TAA GTG ACG	CGA TGG AAG C	CT AAT CGT (CTC TGA ATA	TGG ATC AAA	GAC AGC GTT	TGC TTT TTC 3951
3897	3906	3915	3924	3933	3942	
GGG GCT TCT AGT	TTC TAC AGC G	GAT TTG TGT (GGA TTG TTT	CTT GTT CTG	TTT CTT GGT	TCT TTC TTT 4017
3963	3972	3981	3990	3999	4008	
CTT TTT TTT GTG	TCT CTG TCT G	GCC TTT GTA (CTG CAT GCA	AAC GTG CAC	TCT GAA TGA	TGA ACG ACA
4029	4038		4056	4065	4074	4083
CCA TTT GAC GAT	TGG ATA AGA G	GAT GAC AGA (CTG CAG ATA	CTA TCA TGC	GCA ATG GAA	AAC ACG AAC
4095	4104	4113	4122	4131	4140	4149
AAC CAA GGT TTT	TGA TTC CTT C	AA TAG CGA A	AAT ATA GAA	AAA GAA ACA	AAA AAA AAA	ACA ACA ACA
4161	4170		4188	4197	4206	4215
AAT AAT GGA AGT	ATG ATT AAA C	AC ATT GAG (CGC GAT GAC	TGA CTG GTG	TTG TGA ATG	GCG TGT TGG
4227	4236		4254	4263	4272	4281
TTT TCT TCT TTC 4293	TTG AAA ATT T 4302	AG AAC CGT A	AAA TGT TAT 4320	ATC ATG TGA 4329	TGT AAT GTA 4338	ATA ACA TAT 4347
TTA TAT CTC GTT 4359	GTA TTC TTG T 4368	AC ACA CTT 1	TCC AGG ATA 4386	ACA TGG TCT 4395	GAC ATG GTA 4404	TTT CTG ACG
TAC AAA AAA GAA 4425	AAA GAA AAA C 4434	AG GAA ACC A	ATG AAC CCG 4452	CGA CAA AGC 4461	TGT TCC AGT 4470	TGT TAC AAT 4479
GAT GAT GAT 4491	GAT GAC CTA C	TA CCT AAG (GTA TTC TAT	CTT AGC CAA	GGT ATT CTC	TCG CAT CCT
	4500	4509	4518	4527	4536	4545
ATT CCA TCC TAT	CCT AAC CCG A	GC CTA ACC (CGA GCC TAA	ATA CCT AAA	CTC CTA AAC	TCC TTA ACT 4611
4557	4566	4575	4584	4593	4602	
CCT TAA CTC CTT 4623	TCT AAA TGT C 4632	TA AAC CCC (CAA ACT ATG 4650	AGA CGA CCC 4659	GAA CCC GAA 4668	ACC CTA ATA 4677
AAA GTA TTT ATA	AAC CAT CAT A	AA AGA AAA A	AAA ACC ATC	ATA CAT GGA	TGA TCA AAA	CAA ACA GAA
4689	4698	4707	4716	4725	4734	4743
ACG GAA ACA ACA	CAA CCA GCT A	CC CGC TCA A	AGA CTT TCA	TTC GTT AAT	TCA TCA CTC	ACT CAC TCA
4755	4764		4782	4791	4800	4809
CTC ACT CAC TCA	GCA GCA AAA T	AC CGT TTT C	GTC CTG CTA	TTC GTT TGT	TGC GCC TTG	ATT TCA GGC
4821	4830	4839	4848	4857	4866	4875
GGG ACA ATG GTG 4887	TGA TGT ACG A	CG TGG GGG C 4905	CGG TAG ACT 4914	GCG TCT ACT 4923	GGT GGC ATC 4932	CTT TAC AAT 4941
TTT TTA GTG TGT	CAG TAT GTG A	TG TAT TCA A	ATG CTA TTG	AAC TGA GGG	GGG CTG ATG	GAT AGT GGG
4953	4962	4971	4980	4989	4998	5007
GAG AGA ACA CCT	GAC GGA TAG A	.GG GAA GGA <i>1</i>	ACT GGA CGC	CTG GGG GGA	AGT GAG AGA	GGG GGA TGG
5019		5037	5046	5055	5064	5073
TGG GGA ATA GAT	GAA AAG AGA A	GA GGA GTG A	AGA GCA CAA	GAA GAA AGA	ATG AAT GTT	GGT GAC AAA
5085	5094	5103	5112	5121	5130	5139

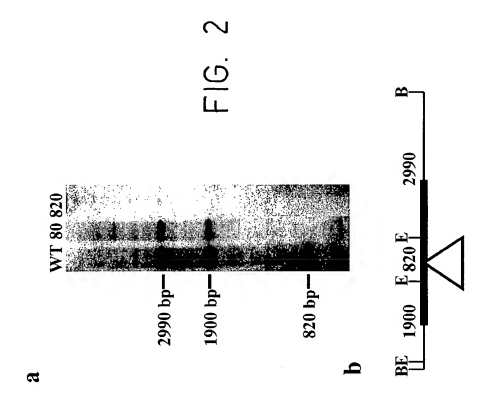
FIG. 1-4

5/7

GTT	AAA GAA 5151	AAG	GAA GGG 5160	GGG	AAA GAG 5169	AAG	AGG	ACA 5178	GGT		GTG 5187		TTG 5196	AGT	AGG 5205	AAG
GGA	AAA AAC 5217	GGA	GAA GGA 5226	AAA	AAA AAA 5235	CAT	AAA	AAA 5244	AAA	AAA	AAA 5253	AAC	AAG 262	AAA	CTA 5271	ACC
AAT	CAT CCA 5283	AAC	TCA GCG 5292	GAA	AGT ACT 5301	CAT	ACA	AAA 5310	GGT		СТG 5319	CCT	TCG 328	GAC	CCA 5337	CAT
TCT	5349	GGT	ACT GAT 5358	TCT	GCT GCC 5367	CCA	GAC	ттс 5376	CAC		CAA 5385	AGT	TAT 394	CAC	ТАТ 5403	TGT
TGT	TAG AGT 5415	GAG	TAG TAG 5424	ACG	TAA GTC 5433	CTC	CCG	ATC 5442	CGG	AGC	CAA 5451	AAC	TCC 460	СТТ	CAG 5469	CTG
TAT	CCC TCT 5481	TCA	ATC CAC 5490	CAG	TAG CAA 5499	CAC	CCA	ТСТ 5508	TGC	CAT 5	AGA 517	GCG	TAT 526	ccc	CCC 5535	CTG
ccc	CTG CCG 5547	AGC	CAG GAG 5556	TAG	CAG TCC 5565	TAT	TCA	TAG 5574	GCG	GAC 5	TCC 5583	TCT	CGT 592	CTT	ACA 601	GGG
ACA	5613	TTG	GTA GGG 5622	CAC	CCG CAG 5631	CAG	AGG	AGG 5640	AGG	ТАТ 5	TTC 649	TGT	GAC 658	TGG	TGT 667	TTG
GGG	CAG CTA 5679	AGG	GCG TGG 5688	GTT	TCC TTC 5697	GTG	AGC 5	CGC 5706	TGT	TGT 5	GAT 715	TGT	CGG 724	CGG	CCG .	AGG
ATA	AGG ATC 5745	С														

FIG. 1-5

6/7



7/7

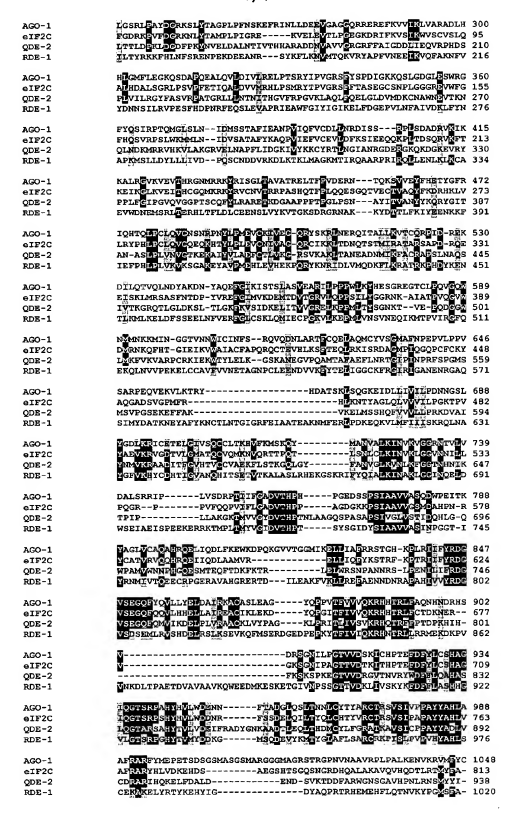


FIG. 3

SEQUENCE LISTING

<110> Università degli Studi di Roma La Sapienza Cogoni, Carlo Macino, Giuseppe Catalanotto, Caterina Azzalin, Gianluca

<120> Isolation and characterization of a N. crassa silencing gene and uses thereof

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<150> RM2000A000021

<151> 2000-01-17

<160> 2

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gtt	gct	ctac	tcg	acaat	tct	gtta	ccaco	ca a	cact	acaa	g tt:	taaca	agtc	atg	ctgaca	540
ato	gtg	gcgg	tcg	tgga	ggt (cgtg	gcggd	g gi	tggt	cgcg	g cg	gcgg	cggc	ggc	ggggag	600
gco	gtg	gagg	tgg	tcago	caa (ggcgg	gcggt	g ga	aggc	cgtg	g ago	gtggt	tac	caaq	gcagcg	660
gcg	igeg	gtgg	agge	ccgt	ggc (ggcgg	gttat	c aa	aggc	ggtgg	g egg	geggt	gac	cgt	gaggcc	720
gtg	ıgcgg	gegg	ttai	tcaaç	igc d	ggtgg	gtggc	g gt	ggtt	tcca	a agg	gegge	ggt	ggaa	ıggggtg	780
gcc	gtgg	gcgg	cggt	ttcc	aa q	ggcgg	gegge	g go	ggcg	gccg	ı tgg	rtggd	ettc	ggcg	gaggac	840
agg	gege	ggg	agga	atacg	aa c	cccc	tcca	ငင္	gato	ıtcta	caa	gtag	ıgtg	cctc	tccatt	900
ttt	tttt	acc	atto	caaca	itg a	tgct	gaca	c ga	cttt	aggg	gaa	ttga	cgg	tcgt	ggtgcc	960
ccc	gago	ctg	acgo	eccag	at c	acca	aact	c ga	ggat	gatt	gga	tcaa	gaa	gcac	gtcagc	1,020
gac	aato	tgg	tcac	ttcc		Ser	aag Lys			Leu					aaa Lys	1071
				Pro			cct Pro									1119
			Trp				ttc Phe 35									1167
							gtt Val									1215
aag Lys 60	gaa Glu	gct Ala	gag Glu	gtc Val	gca Ala 65	tcc Ser	aag Lys	aaa Lys	gtg Val	gag Glu 70	gtg Val	gtg Val	gtt Val	ggg Gly	aaa Lys 75	1263
							aac Asn									1311
							acg Thr									1359
aac	cac	atc	ttt	gag	ata	aco	taa	acc	gag	cca	art	tcc	220	C2.2	226	1407

Asn	Arg	Ile 110		Glu	Val	Thr	Trp		Glu	Pro	Ser	Ser 120		Gln	Asn	
							Trp			aag Lys						1455
										gag Glu 150						1503
										aat Asn				-	_	1551
										gcc Ala				-		1599
										ggt Gly		_			-	1647
										atc Ile						1695
										tta Leu 230						1743
										ctt Leu		_	_		_	1791
										aat Asn						1839
										aga Arg						1887
										ttc Phe						1935
att	gtt	tat	aaa	aaa	tgt	tac	cgc	acg	ctc	aat	ggc	att	gct	aac	cgt	1983

Ile 300		. Tyr	Lys	. Lys	305		Arg	Thr	Leu	Asn 310		, Ile	: Ala	. Asn	1 · Arg 315	
					Lys					Lys					ccg Pro	2031
				Ile											tgt Cys	2079
															cct	2127
		ccc Pro														2175
		caa Gln														2223
		ggc Gly			Glu											2271
		gtc Val											-			2319
gcg Ala								Cys								2367
cag Gln																2415
ctg Leu 460																2463
gtt (Val '			Arg					Pro								2511
aag a	acg	gta	gag	ccg	cag	gac	ggc	ggg	tgg	ttg	atg	aag	ttt	gtc	aag	2559

Lys	Thr	Val	Glu 495	Pro	Gln	Asp	Gly	Gly 500	Trp	Leu	Met	Lys	Phe 505	Val	Lys	
	-	aga Arg 510				_			_				_	_	_	2607
_		tcc Ser	-	_		-			_		-	_		-		2655
		ttc Phe											-			2703
	-	atg Met		_		_			_	_					_	2751
		aag Lys														2799
		aag Lys 590														2847
		ttt Phe														2895
		aag Lys												_		2943
		aag Lys									-	_				2991
		gcc Ala														3039
		aat Asn 670				Gly		_		-	_	-		-		3087
gtc	ggc	ctg	gtc	tca	acc	atc	gac	caa	cac	ctt	gga	caa	tgg	cct	gca	3135

Val	. Gly 685		ı Val	. Ser	Thr	690		Glr.	His	s Leu	695		Trp	Pro	Ala	
	Val					His					Met				ttt Phe 715	3183
					Thr					Trp					gca Ala	3231
				Leu										Gly	gtc Val	3279
										gac Asp						3327
										ggc Gly						3375
										act Thr 790						3423
										agc Ser						3471
										tat Tyr						3519
										cgc Arg						3567
Val										tat Tyr						3615
gac Asp 860				Gln												3663
gcc	acc	aag	gct	gtc	agt	atc	tgc	ccg	cct	gcg	tac	tat	gcc	gac	ttg	3711

Ala Thr Lys Ala Val Ser Ile Cys Pro Pro Ala Tyr Tyr Ala Asp Leu 880 885 890

gtg tgc gac cgg gcg cgt atc cat cag aag gag ctc ttt gac gcc ctc 3759
Val Cys Asp Arg Ala Arg Ile His Gln Lys Glu Leu Phe Asp Ala Leu
895 900 905

gat gaa aac gat agc gtt aag acc gat gat ttc gca aga tgg ggt aac 3807 Asp Glu Asn Asp Ser Val Lys Thr Asp Asp Phe Ala Arg Trp Gly Asn 910 915 920

tcc ggg gct gtt cat ccc aac ctt agg aac tcc atg tac tat atc

Ser Gly Ala Val His Pro Asn Leu Arg Asn Ser Met Tyr Tyr Ile

925

930

935

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7/12

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<211> 938

<212> PRT

<213> Neurospora crassa

<400> 2

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1 5 10 15

Val Arg Pro Gly His Gly Thr Met Gly Glu Lys Val Lys Leu Trp Ala 20 25 30

Asn Tyr Phe Lys Ile Asn Ile Lys Ser Pro Ala Ile Tyr Arg Tyr Thr 35 40 45

Ile Lys Val Ala Ala Thr Glu Glu Lys Leu Gly Lys Glu Ala Glu Val

50	55	60

- Ala Ser Lys Lys Val Glu Val Val Val Gly Lys Leu Leu Lys Gln Ile
 65 70 75 80
- Glu Ala Asn Val Lys Ser Val Ala Ile Ala Ser Asp Phe Lys Val His
- Leu Val Thr Thr Lys Leu Lys Val Pro Glu Asn Arg Ile Phe Glu
 100 105 110
- Val Thr Trp Thr Glu Pro Ser Ser Asn Gln Asn Leu Pro Ser Lys Pro 115 120 125
- Gln Thr Trp Val Val Lys Val Glu Glu Ser Val Glu Thr Cys Asp Phe 130 135 140
- Gly Lys Val Leu Asn Glu Leu Thr Thr Leu Asp Pro Lys Leu Asp Gly
 145 150 155 160
- Asp Phe Pro Lys Tyr Asn Val Glu Leu Asp Ala Leu Asn Thr Ile Val 165 170 175
- Thr His His Ala Arg Ala Asp Asp Asp Val Ala Val Val Gly Arg Gly 180 185 190
- Arg Phe Phe Ala Ile Gly Asp Asp Leu Ile Glu Gln Val Arg Pro His 195 200 205
- Asp Ser Pro Leu Val Ile Leu Arg Gly Tyr Phe Ala Ser Val Arg Pro 210 215 220
- Ala Thr Gly Arg Leu Leu Asn Thr Asn Ile Thr His Gly Val Phe 225 230 230 240
- Arg Pro Gly Val Lys Leu Ala Gln Leu Phe Gln Glu Leu Gly Leu Asp
 245 250 255
- Val Met Asp Lys Cys Asn Ala Trp Asn Glu Val Thr Lys Asn Gln Leu 260 · 265 270
- Asn Asp Lys Met Arg Arg Val His Lys Val Leu Ala Lys Gly Arg Val 275 280 285
- Glu Leu Asn Ala Pro Phe Leu Ile Asp Gly Lys Ile Val Tyr Lys Lys 290 295 300
- Cys Tyr Arg Thr Leu Asn Gly Ile Ala Asn Arg Gly Asp Glu Arg Gly

305		•		310					315					320
Tue Cle	T	n	C1	T	01	17- 1	7	m	D	D	-	5 1.	6 1	- 1

- Lys Gln Lys Asp Gly Lys Glu Val Arg Tyr Pro Pro Leu Phe Gly Ile 325 330 335
- Pro Gly Val Gln Val Gly Gly Pro Thr Ser Cys Gln Phe Tyr Leu Arg 340 345 350
- Ala Arg Glu Thr Lys Asp Gly Ala Ala Pro Pro Pro Thr Pro Gly Leu 355 360 365
- Pro Ser Asn Ala Tyr Ile Thr Val Ala Asn Tyr Tyr Lys Gln Arg Tyr 370 380
- Gly Ile Thr Ala Asn Ala Ser Leu Pro Leu Val Asn Val Gly Thr Lys 385 390 395 400
- Glu Lys Ala Ile Tyr Val Leu Ala Glu Phe Cys Thr Leu Val Lys Gly
 405 410 415
- Arg Ser Val Lys Ala Lys Leu Thr Ala Asn Glu Ala Asp Asn Met Ile 420 425 430
- Lys Phe Ala Cys Arg Ala Pro Ser Leu Asn Ala Gln Ser Ile Val Thr 435 440 445
- Lys Gly Arg Gln Thr Leu Gly Leu Asp Lys Ser Leu Thr Leu Gly Lys 450 455 460
- Phe Lys Val Ser Ile Asp Lys Glu Leu Ile Thr Val Val Gly Arg Glu 465 470 475 480
- Leu Lys Pro Pro Met Leu Thr Tyr Ser Gly Asn Lys Thr Val Glu Pro 485 490 495
- Gln Asp Gly Gly Trp Leu Met Lys Phe Val Lys Val Ala Arg Pro Cys
 500 505 510
- Arg Lys Ile Glu Lys Trp Thr Tyr Leu Glu Leu Lys Gly Ser Lys Ala 515 520 525
- Asn Glu Gly Val Pro Gln Ala Met Thr Ala Phe Ala Glu Phe Leu Asn 530 535 540
- Arg Thr Gly Ile Pro Ile Asn Pro Arg Phe Ser Pro Gly Met Ser Met 545 550 555 560
- Ser Val Pro Gly Ser Glu Lys Glu Phe Phe Ala Lys Val Lys Glu Leu

565 570 575

Met Ser Ser His Gln Phe Val Val Leu Leu Pro Arg Lys Asp Val 580 585 590

- Ala Ile Tyr Asn Met Val Lys Arg Ala Ala Asp Ile Thr Phe Gly Val 595 600 605
- His Thr Val Cys Cys Val Ala Glu Lys Phe Leu Ser Thr Lys Gly Gln 610 615 620
- Leu Gly Tyr Phe Ala Asn Val Gly Leu Lys Val Asn Leu Lys Phe Gly 625 630 635 640
- Gly Thr Asn His Asn Ile Lys Thr Pro Ile Pro Leu Leu Ala Lys Gly
 645 650 655
- Lys Thr Met Val Val Gly Tyr Asp Val Thr His Pro Thr Asn Leu Ala 660 665 670
- Ala Gly Gln Ser Pro Ala Ser Ala Pro Ser Ile Val Gly Leu Val Ser 675 680 685
- Thr Ile Asp Gln His Leu Gly Gln Trp Pro Ala Met Val Trp Asn Asn 690 695 700
- Pro His Gly Gln Glu Ser Met Thr Glu Gln Phe Thr Asp Lys Phe Lys 705 710 715 720
- Thr Arg Leu Glu Leu Trp Arg Ser Asn Pro Ala Asn Asn Arg Ser Leu 725 730 735
- Pro Glu Asn Ile Leu Ile Phe Arg Asp Gly Val Ser Glu Gly Gln Phe 740 745 750
- Gln Met Val Ile Lys Asp Glu Leu Pro Leu Val Arg Ala Ala Cys Lys 755 760 765
- Leu Val Tyr Pro Ala Gly Lys Leu Pro Arg Ile Thr Leu Ile Val Ser 770 780
- Val Lys Arg His Gln Thr Arg Phe Phe Pro Thr Asp Pro Lys His Ile 785 790 795 800
- His Phe Lys Ser Lys Ser Pro Lys Glu Gly Thr Val Val Asp Arg Gly 805 810 815
- Val Thr Asn Val Arg Tyr Trp Asp Phe Phe Leu Gln Ala His Ala Ser

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820 825 830 Leu Gln Gly Thr Ala Arg Ser Ala His Tyr Thr Val Leu Val Asp Glu 835 840 845 Ile Phe Arg Ala Asp Tyr Gly Asn Lys Ala Ala Asp Thr Leu Glu Gln $^{\circ}$ 855 Leu Thr His Asp Met Cys Tyr Leu Phe Gly Arg Ala Thr Lys Ala Val 870 875 Ser Ile Cys Pro Pro Ala Tyr Tyr Ala Asp Leu Val Cys Asp Arg Ala 885 890 Arg Ile His Gln Lys Glu Leu Phe Asp Ala Leu Asp Glu Asn Asp Ser 900 . 905 Val Lys Thr Asp Asp Phe Ala Arg Trp Gly Asn Ser Gly Ala Val His

920

925

Pro Asn Leu Arg Asn Ser Met Tyr Tyr Ile 930 935

915